

Distribution Patterns of Postmortem Damage in Human Mitochondrial DNA

M. Thomas P. Gilbert,¹ Eske Willerslev,^{2,*} Anders J. Hansen,^{2,*} Ian Barnes,^{1,†} Lars Rudbeck,³ Niels Lynnerup,⁴ and Alan Cooper¹

¹Henry Wellcome Ancient Biomolecules Centre, Department of Zoology, Oxford University, Oxford, United Kingdom; and ²Department of Evolutionary Biology, Zoological Institute, and ³Research Laboratory and ⁴Laboratory of Biological Anthropology, Institute of Forensic Medicine, University of Copenhagen, Copenhagen

The distribution of postmortem damage in mitochondrial DNA retrieved from 37 ancient human DNA samples was analyzed by cloning and was compared with a selection of published animal data. A relative rate of damage (ρ_v) was calculated for nucleotide positions within the human hypervariable region 1 (HVR1) and cytochrome oxidase subunit III genes. A comparison of damaged sites within and between the regions reveals that damage hotspots exist and that, in the HVR1, these correlate with sites known to have high *in vivo* mutation rates. Conversely, HVR1 subregions with known structural function, such as MT5, have lower *in vivo* mutation rates and lower postmortem-damage rates. The postmortem data also identify a possible functional subregion of the HVR1, termed “low-diversity 1,” through the lack of sequence damage. The amount of postmortem damage observed in mitochondrial coding regions was significantly lower than in the HVR1, and, although hotspots were noted, these did not correlate with codon position. Finally, a simple method for the identification of incorrect archaeological haplogroup designations is introduced, on the basis of the observed spectrum of postmortem damage.

Introduction

The extreme susceptibility that research on human ancient DNA (aDNA) has to contamination with modern human DNA has caused skepticism about the authenticity of many results (Richards et al. 1995; Stoneking 1995; Handt et al. 1996; Cooper 1997; Kolman and Tuross 2000; Cooper et al. 2001*b*). Several studies have shown that, despite rigorous protocols (e.g., see Cooper and Poinar 2000), contaminants can still be identified in amplified products (Richards et al. 1995; Handt et al. 1996; Krings et al. 1997; Kolman and Tuross 2000; Hofreiter et al. 2001). Contaminants are usually identified when repeat amplifications yield differing, or multiple, sequences in cloned products. However, the effect that postmortem damage has on endogenous DNA is often underrated. The majority of postmortem DNA damage occurs as double-strand breaks and oxidative dinucleotide modification, both of which prevent sub-

sequent enzymatic replication (Pääbo 1989; Lindahl 1993; Höss et al. 1996). However, minor sequence modifications, such as hydrolytic deamination and depurination, permit polymerase action and are manifested as limited amounts of base variation among sequenced clones (Krings et al. 1997). Few studies have examined the prevalence of such postmortem damage in detail (Handt et al. 1996; Kolman and Tuross 2000; Hansen et al. 2001; Hofreiter et al. 2001), and it is generally considered that, when the initial template number is >1,000 copies, postmortem-damage rates are unlikely to bias results (Handt et al. 1996; Krings et al. 1997). However, when few DNA templates initiate a PCR, the resulting sequences are likely to contain artifacts.

mtDNA is generally present in archaeological samples in much greater amounts than nuclear DNA is (Greenwood et al. 1999), and, consequently, most human ancient genetic analyses have used the well-characterized hypervariable region 1 (HVR1) (e.g., see Oota et al. 1995, 1999; Ribeiro-dos-Santos et al. 1996; Torroni et al. 1998; Lalueza-Fox et al. 2001). Archaeological HVR1 results can be compared with detailed records of base substitutions within modern populations, and sequences can be located within the commonly used human phylogenetic tree, relative to the Cambridge reference sequence (CRS) (Anderson et al. 1981). Unfortunately, a major limitation of this classification system is that certain haplogroups, especially those of Eurasian origin, are routinely categorized using fewer than five site changes—and often as few as one. If these key

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Address for correspondence and reprints: Dr. Alan Cooper, Henry Wellcome Ancient Biomolecules Centre, Department of Zoology, University of Oxford, South Parks Road, Oxford OX1 3PS, United Kingdom. E-mail: alan.cooper@zoo.ox.ac.uk

* The second and third authors contributed equally to this work.

† Present affiliation: Department of Biology, Darwin Building, University College London, London.

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sites are commonly modified by postmortem damage, then haplogroup designations may be incorrect. In one of the only studies of this issue, Hofreiter et al. (2001) suggested that postmortem damage appeared to be randomly distributed in the control region of the cave bear (*Ursus spelaeus*). However, only a small number of samples were studied, and it is also possible that postmortem damage may vary according to environmental or taxonomic factors.

Within modern human populations, certain nucleotide positions within the HVR1, termed “sites,” appear to mutate at significantly higher rates than others, and these sites have been termed “hotspots” (Hasegawa and Horai 1991; Vigilant et al. 1991; Hasegawa et al. 1993; Wakely 1993; Aris-Brisou and Excoffier 1996; Macauley et al. 1997; Excoffier and Yang 1999; Meyer et al. 1999; Stoneking 2000; Finnillä et al. 2001; Heyer et al. 2001). It has been suggested that the variation in rates of damage is related to DNA secondary and tertiary structure (Heyer et al. 2001), and it is possible that the same sites might also be susceptible to postmortem damage. To investigate these issues, we amplified and cloned human HVR1 and cytochrome oxidase subunit III (COIII) mtDNA from a range of archaeological specimens, and we compared the human sequences to aDNA sequences from brown bear (*U. arctos*), domestic cow (*Bos taurus*), pig (*Sus scrofa*), goat (*Capra hircus*), and sheep (*Ovis aries*), as well as to published data from other taxa.

Material and Methods

Samples

It is exceedingly difficult to authenticate human aDNA sequences, and it may even be impossible in certain cases (Cooper 1997). Several authors have demonstrated how even stringent controls can fail to prevent or detect contamination (Handt et al. 1996; Kolman and Tuross 2000). Consequently, careful attention was paid to the choice of methods and samples, to provide the best chance of limiting contamination. Forty-three whole teeth were obtained from 34 archaeological human skeletons preserved in a range of temporal, geographical, and environmental situations (table 1). Teeth were used as a DNA source both because the relatively impervious outer enamel layer provides a degree of protection from contaminating DNA sources (M.T.P.G., unpublished data) and because teeth have been shown to yield higher amounts of DNA than bone does in many environments (Kurosaki et al. 1993; Oota et al. 1995). Teeth from cow, pig, goat, and sheep specimens obtained from the Greenland archaeological site were also analyzed for human DNA as a contamination control. Multiple teeth were taken from five specimens, to allow replication of the extraction and amplification procedures. DNA from one of

these five specimens was extracted and amplified independently at the Ancient DNA Laboratory (Zoological Institute, Copenhagen) by using similar techniques, to confirm that the spectrum of sequence damage observed was replicable (tooth EA). Two brown bear 12s mtDNA sequences were also analyzed, to allow taxonomic comparisons of sequence damage outside the HVR1. Both bear sequences have previously been replicated at the University of California–Los Angeles and Oxford University (Barnes et al. 2002).

Extraction

All DNA-preparation and -extraction methods followed strict aDNA-specific requirements (Cooper and Poinar 2000) and were performed in a dedicated aDNA facility, in a building physically isolated from any molecular biology research. Full-body suits, breathing masks, and face shields were used, and gloves were frequently changed. DNA was extracted from teeth by using a new technique that significantly reduces external contamination (fig. 1) (M.T.P.G., unpublished data). Whole teeth were initially washed in 50% bleach for 5 min, followed by exposure to 254-nm-wavelength UV light for 10 min on either side. Teeth were fully encased, upside down, in RTV-11 liquid silicone rubber (Tiranti) and were left overnight while the matrix hardened. The silicone that encases the top 5–10 mm of the root was removed with a horizontal cut, and the root was removed flush with the silicon by using a single-use carborundum cutting disk. Dental pulp was powdered and removed from the pulp chamber by using a dental drill bit and was digested and extracted using a phenol:chloroform DNA-extraction protocol (after Barnes et al. 2002), with 0.2 M PTB (N-phenacylthiazolium bromide) (Poinar et al. 1998). PTB has been shown to increase DNA yield in aDNA extractions, putatively by cleaving cross-linked proteins and DNA formed by condensation reactions (Vasan et al. 1996; Poinar et al. 1998; Bada 1999). The bear DNA extractions were as described elsewhere (Barnes et al. 2002), except with the addition of 0.001 M PTB.

PCR Amplification

PCR amplifications used a high-fidelity polymerase (Platinum *Taq* Hi-Fidelity; Invitrogen) (after Cooper et al. 2001a) to reduce the polymerase-error rate and to increase amplification efficiency (Willerslev et al. 1999; Hansen et al. 2001; Gilbert et al. 2003 [in this issue]). Full details of the mitochondrial regions amplified are given in table 1, and primers and annealing temperatures are given in table 2. All samples were amplified and extensively cloned at least twice. PCR products were purified, by precipitation, using Microclean (Microzone) and were cloned using the Topo TA cloning system (Invitrogen). Colonies were used to initiate PCR

Table 1

Details of Samples Studied

SPECIES AND SAMPLE	SITE ^a	SAMPLE DATE	HVR1				COIII ^b			
			Amps ^c	Clones ^d	UNG ^e	UNG Clones ^f	Amps ^c	Clones ^d	UNG ^e	UNG Clones ^f
<i>Homo sapiens:</i>										
Tg44	Greenland	1000–1200 A.D. ^g	1	2	1	8				
Tg54	Greenland	1000–1200 A.D. ^g	1	5	1	8				
Tg63 ^h	Britain	Not known	1	32	1	12				
Tg76	Denmark	1711 A.D. ⁱ	1	5	1	11	1	12	1	12
Tg77 (a) ^j	Denmark	1400–1500 A.D. ^k	3	9	1	12				
Tg80	Denmark	1300–1600 A.D. ^l	1	7	1	11	1	10	1	9
Tg85	Denmark	1711 A.D. ⁱ	1	5	1	12				
Tg99.1 (e) ^{h,j}	Greenland	1000–1200 A.D. ^g	9	60	1	20				
Tg99.2 (e) ^j	Greenland	1000–1200 A.D. ^g	1	16						
Tg99.3 (e) ^j	Greenland	1000–1200 A.D. ^g	1	16						
Tg103	Denmark	1300–1600 A.D. ^l	1	4		11				
Tg104	Britain	Not known	1	15	1	16				
Tg105	Denmark	400–1000 A.D. ^l	1	4	1	6				
Tg112	Greenland	1000–1200 A.D. ^g	1	6	1	7				
Tg114	Denmark	1711 A.D. ⁱ	1	5	1	12	1	11	1	12
Tg116	Denmark	700–1000 A.D. ^l	2	10	1	12				
Tg120	Denmark	1711 A.D. ⁱ	1	2	1	8				
Tg123	Denmark	1400–1500 A.D. ^k	1	6	1	8				
Tg127 (a) ^j	Denmark	1711 A.D. ⁱ	1	8	1	11				
Tg128	Greenland	1000–1200 A.D. ^g	2	12						
Tg129	Greenland	1000–1200 A.D. ^g	2	9						
Tg131	Greenland	1000–1200 A.D. ^g	1	7	1	7				
Tg133	Greenland	1000–1200 A.D. ^g	1	2	1	8				
Tg136.1	Britain	1172–1536 A.D.	2	17			1	11	1	12
Tg136.2	Britain	1172–1536 A.D.	2	26						
Tg137.1 (c) ^j	Britain	675–740 A.D.	3	27						
Tg137.2 (c) ^j	Britain	675–740 A.D.	3	38						
Tg138 (b) ^j	Britain	873–874 A.D.	1	8	1	5				
Tg141	Britain	873–874 A.D.	1	6	1	6				
Tg142	Britain	873–874 A.D.	1	7	1	9				
Tg143	Britain	874–876 A.D.	1	6	1	8				
Tg145	Britain	750–873 A.D.	1	5	1	7				
Tg146	Britain	900–1100 A.D.	1	6	1	8				
Tg147 ^h	Britain	873–874 A.D.	2	9						
Tg148 (c) ^j	Britain	675–740 A.D.	7	80	1	21	3	33	1	12
Tg149 (b) ^j	Britain	873–874 A.D.	7	57	1	12	2	21	1	10
Tg191 ^h	Britain	Not known	2	8						
Tg192	Britain	Not known	2	10	1	11	1	12	1	12
Tg196	Not known	Not known	2	12						
Tg232 (d) ^j	Britain	Not known	1	8	1	8				
Tg233 (d) ^j	Britain	Not known	2	14						
Tg200 (e) ^{h,j}	Greenland	1000–1200 A.D. ^g	1	15	2	19				
EA (e) ^j	Greenland	1000–1200 A.D. ^g	1							
<i>U. arctos:</i>										
IB221	Alaska	Not known					1	12		
IB223	Alaska	Not known					1	12		
Cow (<i>B. taurus</i>)	Greenland	1000–1200 A.D. ^g	1 ^m							
Sheep (<i>O. aries</i>)	Greenland	1000–1200 A.D. ^g	1 ^m							
Pig (<i>S. scrofa</i>)	Greenland	1000–1200 A.D. ^g	1 ^m							
Goat (<i>C. hircus</i>)	Greenland	1000–1200 A.D. ^g	1 ^m							
Total			83	606	31	314	12	134	7	79

^a Greenland samples have been identified as Viking (Lynnnerup 1998).^b The 12s gene was amplified in bear samples.^c Number of untreated (i.e., no UNG) PCR per sample.^d Total number of non-UNG clones sequenced per sample.^e Number of PCR with UNG treatment per sample.^f Total number of UNG clones sequenced per sample.^g Dated on the basis of radiocarbon and archaeological analysis (Arneborg et al. 1999).^h Sample rejected owing to signs of contamination.ⁱ Dated on the basis of historical records (Copenhagen City Museum, Copenhagen, Denmark).^j Multiple teeth originating from one of five skeletons (labeled, in parentheses, as “a”–“e”).^k Dated on the basis of archaeological analysis (Aalborg Museum, Aalborg, Denmark).^l Dated on the basis of archaeological analysis (Panum Institute, Copenhagen, Denmark).^m No PCR product.

reamplifications with vector M13R and T7 primers, were purified as before, and were sequenced on an ABI 377 or 3700 by using the ABI Big Dye 3 PRISM kit. Blank control extractions were performed at a ratio of 1:5 samples, and negative control amplifications were performed at a ratio of 1:1. No positive controls were used at any step of the amplification process.

Uracil-N-Glycosylase (UNG) Treatment

A 5-μl aliquot of each human DNA extract was treated with 1 U *Escherichia coli* UNG (Sigma), to excise uracil caused by hydrolytic deamination of cytosine (Dinner et al. 2001). UNG reduces sequence artifacts caused by this common form of postmortem damage, resulting in an apparent C→T/G→A mutation (Pääbo 1989; Hofreiter et al. 2001). After UNG treatment, extracts containing multiple sources of DNA can be identified as those in which cloned sequences still either share no consensus sequence or retain sporadic C→T/G→A base changes that previously may have been attributed to postmortem damage but that now may be recognized as authentic base differences between a contaminant and the sample's endogenous sequence.

Damage Spectrum in Ancient Specimens

To examine the robustness of the results, we examined samples from a range of sites and time periods (table 1). Previously published cloned aDNA sequences of cave bear (Loreille et al. 2001) and brown bear (Barnes et al. 2002) were obtained (see table 3). Other reported human aDNA clones cannot be included in the present analysis because of an inability to determine the extent of contamination in those studies.

Contamination

Contaminating DNA is a major concern, because it can imitate sequence damage and also permits jumping PCR between endogenous and contaminant strands (Pääbo

Table 2

Details of Primer Used

Primer	Sequence	Annealing Temperature (°C)
L09486	5' ttc gca gga ttt ttc tga gc	56
H09652	5' tgg tga gct cag gtg att ga	56
12se	Barnes et al. 2002	
12sf3	Barnes et al. 2002	
L16055	Handt et al. 1996	
H16410	Handt et al. 1996	
L16209	Handt et al. 1996	
H16356	Handt et al. 1996	

NOTE.—Conditions for previously reported primers were identical to the references.

et al. 1990). Jumping events will increase the apparent number of damaged sites in cloned sequences by introducing positions that differ between the contaminant and authentic DNA. Low-level contamination often is neither observed nor reported, but this insidious problem would be manifested in clone sequences as sporadic, apparently postmortem damage at phylogenetically variable sites. So that this problem may be avoided, we analyzed specimens representing a variety of mitochondrial haplogroups, to detect common contaminants, and used at least two separate PCR amplifications and UNG treatment (which involves a further PCR replication), to confirm that extracts contain only one source of ancient sequence. It could be argued that an old—and therefore damaged—contaminant could generate a solitary consensus sequence from a tooth and that UNG treatment would confirm that the observed sequence heterogeneity had arisen from postmortem damage (as opposed to several contaminating sequences). Although this situation appears to be rare, such results would not affect the analyses, since DNA damage is the only issue under investigation, whether it is from ancient endogenous or contaminant sources.

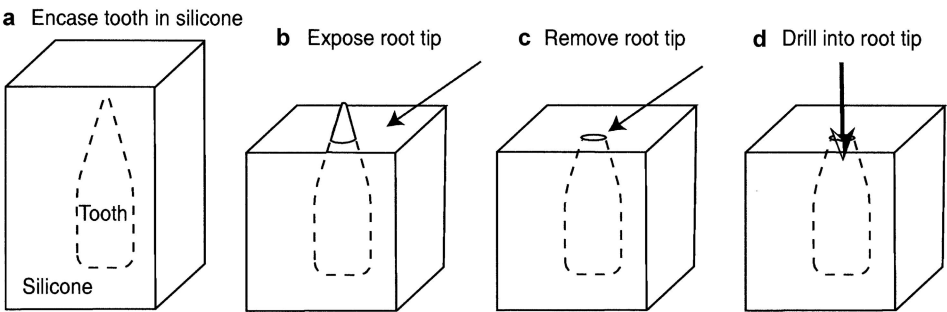


Figure 1 Powder-extraction method. *a*, Encasement of tooth, crown down, in liquid silicone, which is left to set. *b*, Exposure of root tip (5–10 mm) by using horizontal cuts into the silicone. *c*, Horizontal removal of root tip, flush against silicone surface, with sterile cutting disk. *d*, Extraction of sterile pulp-cavity powder by drilling into root from cut surface.

Table 3

Haplogroups Ranked by Misidentification Potential in Using a Predominantly Haplogroup H and Haplogroup V Data Set

Haplogroup and Site	Site Rate ^a	Lowest Rate ^b
U5:		15.4
270	15.4	
B:		12.5†
189	12.5†	
J:		12.5†
126	12.5†	
69	12.5†	
JT*:		12.5†
126	12.5†	
D:		10.3†
362	12.5†	
223	10.3†	
G:		10.3
223	10.3	
L3a*:		10.3
223	10.3	
M*:		10.3
223	10.3	
N*:		10.3
223	10.3	
E:		10.3
223	10.3	
A:		6.4
319	6.4	
290	7.7	
223	10.3	
C:		6.4
327	7.7	
298	6.4	
223	10.3	
Pre-C:		6.4
298	6.4	
223	10.3	
T:		6.4
294	6.4	
126	12.5†	
V:		6.4
298	6.4	
X:		6.4
278	6.4	
223	10.3	
F:		3.8
304	3.8	
M1:		2.6
311	6.4	
249	2.6	
223	10.3	
189	12.5†	
129	12.5†	
U1:		2.6
249	2.6	
189	12.5†	
W:		2.6
292	2.6	
223	10.3	
K:		2.6
311	6.4	
224	2.6	
L3a1:		1.3
320	1.3	
223	10.3	
I:		.0†
391	.0†	
223	10.3	
129	12.5†	

(continued)

Table 3 (continued)

Haplogroup and Site	Site Rate ^a	Lowest Rate ^b
L3a2:		.0†
223	10.3	
209	.0†	
M2:		.0
343	.0	
311	6.4	
265	1.3	
241	3.8	
223	10.3	
148	.0†	
144	37.5†	
129	12.5†	
N1:		.0
223	10.3	
176	.0†	
145	.0†	
N2:		.0
248	7.7	
223	10.3	
147	.0†	
N3:		.0
261	.0	
257	2.6	
223	10.3	
U3:		.0
343	.0	
U4:		.0
356	.0	
U6:		.0†
219	1.3	
172	.0†	

NOTE.—Rates with a dagger (†) following are estimates on the basis of the OR, and rates without are estimates on the basis of the MR. For full details, see the “Haplogroup Analysis” subsection.

^a Relative rate per site.

^b Haplogroup minimum value.

Determination of Site-Specific Postmortem-Damage Rates

Bases at CRS positions 16209–16356 (here termed the “middle region” [MR]) contain the majority of phylogenetically variable sites and were consequently subject to more amplifications than were the outer regions (OR), at positions 16055–16208 and 16357–16410. Four samples that produced multiple sequences in every cloned product were presumed to be contaminated and were removed from the analysis. Cloned sequences from the remaining samples were aligned manually with the CRS by using the program Sequence Navigator (version 1.0.1; Applied Biosystems). The consensus sequence for each specimen was determined from the sequences shared between all clones, including the UNG-treated amplifications. The remaining intracclone base differences formed the postmortem-damage data set. The few base insertions and deletions observed were excluded from the analysis.

It is difficult to calculate site-specific HVR1 postmortem-damage rates for a number of reasons. A ma-

major problem is that damaged sites occurring on several clones within a set derived from one PCR are likely to have arisen from the same ancestral template molecule and, therefore, that the observed number is dependent on the number of clones sequenced. Similarly, the proportion of undamaged starting templates is difficult to assess because of replication advantages over damaged templates during the early amplification cycles. Furthermore, within a set of clones, it is common to see jumping PCR spread damaged sites between daughter amplified strands, generating a few damaged sites at identical nucleotide positions between quite different sequences. Last, the different number of PCR amplifications and clones for each fragment (MR and OR) complicate calculations. Consequently, an approximate relative rate of postmortem damage (ρ_v) was calculated for each site after the method of Hofreiter et al. (2001), which was modified to account for the dual-fragment amplification of the HVR1 (fig. 2). A modified relative rate of postmortem damage was calculated as $\rho_v = \mu_v/\sigma_v$, where v is a specific site with reference to the CRS, μ_v is the number of hits observed at a specific site across all sequences analyzed, and σ_v is the total number of amplifications for each specific site. This calculation is biased, since it does not take into account the numbers of clones examined at each site (which vary by PCR) or the problems associated with assessing the number of undamaged starting templates. Some previous studies (e.g., see Cooper et al. 2001a) have assumed that the low number of starting templates implies that all undamaged strands effectively arose from one template. The denominator then becomes the number of different clones (as opposed to PCRs). However, our data demonstrate that different genetic regions exhibit heterogeneity in the rates of postmortem damage and, therefore, that an area with larger amounts of background postmortem damage will have a higher denominator in the rate calculation, lowering the apparent relative rates of damage for sites in this region. Consequently, it is difficult to compare postmortem-damage rates between different subregions of the HVR1 or between the HVR1 and COIII, and such calculations should be seen as illustrative, not absolute. It is, however, possible to compare postmortem-damage estimates within a single region of amplification (MR or OR), since all sites have been amplified/cloned the same number of times.

Hotspots

The null hypothesis, H_0 , that postmortem-damaged sites were randomly distributed across the HVR1 and COIII was tested through comparison with the expected Poisson distribution (Aris-Brisou and Excoffier 1996; Heyer et al. 2001). Heyer et al. (2001) have calculated

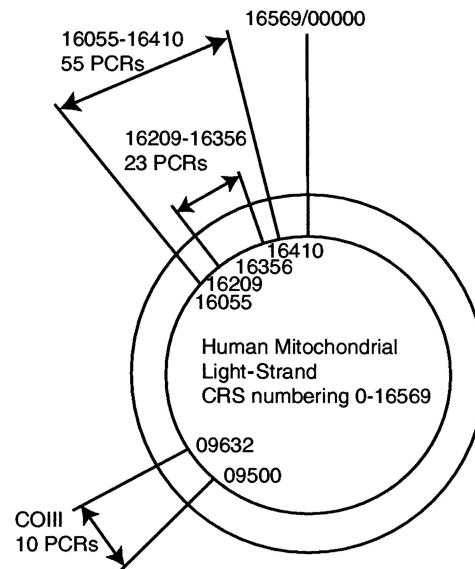


Figure 2 Human mitochondrial regions amplified

an expected distribution of substitutions in modern human mitochondrial samples and have tested this against empirical data. It is a simple matter to modify the test for this data set, although, because of the bias described above (see the “Determination of Site-Specific Postmortem-Damage Rates” subsection), the HVR1 analysis was performed separately on the MR and the OR. The probability that sequence sites have exactly X substitutions, $P(X)$, is $P(X) = e^{-\lambda}\lambda^X/X!$, where Poisson parameter λ refers to the observed density of mutations. To estimate the expected count, $LP(X)$, of sequence positions for each category, given the random distribution λ and the length L of the sequence, we multiply the Poisson probability $P(X)$ by the number of sequence sites L , such that $LP(X) = L(e^{-\lambda}\lambda^X/X!)$. A χ^2 goodness-of-fit test can then be applied to the observed and expected results, to determine whether the H_0 that postmortem damage is randomly distributed within each of the regions can be rejected.

Relative-Rate Comparison with Modern Studies

To determine whether there is a correlation between sites with elevated substitution rates in modern populations and sites with high postmortem-damage rates, we obtained estimates of site-specific rates of mutation from recent comprehensive studies (Excoffier and Yang 1999; Meyer et al. 1999). Different rate-calculation methods were used in each study, preventing direct comparison, so the results were standardized onto a quartile scale of 1–4 (with 1 being the lowest and 4 being the highest), excluding invariant sites. Sites with consistently

high mutation rates (3 or 4) in both modern studies were contrasted with those in the postmortem data set.

Base Structure

The originally damaged mtDNA strand (light or heavy) was determined according to the method described in the companion article by Gilbert et al. (2003), by using the prevalence of hydrolytic deamination of cytosine to deoxycytidine residues (leading to an observable C→T or G→A change; C→T/G→A), which is by far the most commonly observed form of postmortem damage (Hansen et al. 2001; Hofreiter et al. 2001). The base composition around the most mutable and damaged HVR1 sites (those with a standardized mutation/damage rate of 3 or 4) were examined for patterns by using a window of 3, 5, 7, 9, or 11 bp around the hotspot. Principal-component analysis was performed on the five data sets, and the H_0 that the first and second eigenvectors have no correlation with levels of postmortem damage was tested using a general linear model.

Functional Analysis

The rate of postmortem damage and variation in modern data sets was analyzed in three short regions of known function within the OR: 7sDNA, MT5, and TAS (positions 16106–16191, 16194–16208, and 16157–16172, respectively) (Doda et al. 1981; Ohno et al. 1991). An additional area of low variation, observed in the postmortem data set (low-damage region 1 [LDR1]) at positions 16365–16395, was also compared. The H_0 that the percentage of variable bases did not differ significantly along the OR was tested using a Student's unpaired two-sample t test and a Mann-Whitney test. For the COIII postmortem data set, the H_0 that postmortem damage (measured as total hits per codon position [hereafter "hits"]) and the number of variable sites at each codon position (hereafter "sites") were evenly distributed among different codon positions was tested by comparing the observed and expected levels with a χ^2 goodness-of-fit test.

Regional Specificity

Postmortem-damage rates in the HVR1 and COIII can be compared if it is assumed that amplifications from the same DNA extract will (a) start from the same template number and combination of postmortem-damaged and undamaged strands and (b) amplify under similar conditions. A comparison of relative rates can be made by using the number of clones screened for HVR1 and COIII and assuming that the number of undamaged sequences results from the same relative number of undamaged templates. The calculation of comparable rate, R , is $R = T/L\Pi$, where T is the total number of hits, L is the total length of sequence, and Π is the number of

PCR amplifications. A saturation value, S , was also calculated for each sample, representing the number of different sequences within the clones. An S value of 1 indicates that all sequences are different, and, in such cases, it is probable that further clones would yield different products and, therefore, that postmortem damage for the individual has been underestimated. The H_0 that no difference exists in the proportion of sites that show damage between HVR1 and COIII for each sample was tested using a Student's one-sample t test and a Wilcoxon test.

Haplogroup Analysis

For the analysis of the possible phylogenetic effects of damage-induced sequence artifacts, a Eurasian mitochondrial tree based predominantly on HVR1 sequences (Richards and Macaulay 2000) was modified to show which branches were defined by sites with high postmortem-damage rates. This comparison requires that the strand (light or heavy) on which the postmortem damage occurred be identified, to allow determination of the direction in which the haplogroup assignments will shift on the phylogenetic tree. For the standardization of the postmortem-damage rates between the OR and the MR, the OR rates were multiplied by the ratio of the number of PCRs over the MR and the OR.

Results

Data Authenticity, Postmortem Damage, and UNG

Of the original 43 human teeth samples, 4 (Tg99.1, Tg147, Tg191, and Tg200) appeared to be contaminated, since cloned sequences from a single PCR amplification differed significantly and/or repeat amplifications yielded different products. Tg99.1 contained at least six sequences, whereas Tg147, Tg191, and Tg200 contained two different sequences each. All other samples possessed the same consensus sequence in the separate amplification and cloning experiments. Of the 34 sampled individuals, 7 shared the haplogroup (V) of the researcher (M.T.P.G.) undertaking the experimental analysis (M.T.P.G., unpublished data). However, the sporadic base changes among the clones suggest that these sequences are ancient. All other samples used for the analysis (including the sample sent for independent replication) displayed similar postmortem hydrolytic damage. The four animal teeth examined, from two of the archaeological sites, repeatedly yielded no human DNA, and tests using "spiked" PCR revealed that this did not result from PCR inhibition.

A consistent, single haplotype may easily be obtained when a sample with no endogenous DNA is contaminated by a single modern source or even several sources if one is more concentrated than the others (Cooper

1997). However, cloned sequences of all the postmortem samples show a pattern of a shared consensus sequence with many scattered singleton substitutions, particularly C→T/G→A, characteristic of aDNA (Handt et al. 1996; Krings et al. 1997; Hansen et al. 2001; Gilbert et al. 2003). In samples treated with UNG, no C→T/G→A changes were observed except where shared between all clones (therefore deemed to be part of the original sequence). This strongly suggests that the sequences derive from one ancient source (a damaged contaminant that is either endogenous or old) and are therefore suitable for the present study. The UNG results, together with the very low rate of polymerase misincorporation observed in modern contaminants from both modern DNA and aDNA extractions (Gilbert et al. 2003), indicate that the observed sequence heterogeneity in ancient sequences is not a result of enzyme misincorporation.

Rates of Postmortem Damage

The spatial variation of rates of postmortem damage per site—calculated as absolute number of hits, A_n , for COIII, and as A_n in addition to the more comparable relative rate, ρ_n , of postmortem damage, for the HVR1—can be seen in figure 3. The variation initially appears to be overdispersed, with certain sites being overrepresented, such as sites 16110, 16204, 16223, 16270, 16298, and 16325, in the HVR1, and sites 9540, 9545, and 9570, in the COIII. The H_0 that postmortem damage is randomly distributed can be strongly rejected (fig. 4) (MR $\chi^2 P < .00$; OR $P = .02$; COIII $P < .00$), demonstrating that sites are not being damaged at random in either region.

Structural Analysis

The majority (24/30) of sites with a high rate of postmortem damage have also been observed as hypermutable in the modern data sets (table 4). The exceptions (sites 16110, 16131, 16144, 16204, 16242, and 16325) are of special interest, and the surrounding base composition was examined for a possible cause. The first and second eigenvectors of the principal-component analysis explained the majority of the results in all cases, and there was no evidence to reject the H_0 that base composition around individual sites plays no role in determining a propensity for postmortem damage (results not shown). However, two of the three HVR1 subregions with known function appear to have a reduced rate of postmortem damage (fig. 4b). A fourth region of low postmortem damage, LDR1, is also apparent at sites 16365–16395 in the HVR1. Two-sample t tests and non-parametric Mann-Whitney tests show that the percentage of postmortem-damaged sites within LDR1 is significantly different from the average rate across the OR,

at $P < .05$, and across MT5, at $P < .1$ (table 5). Within COIII, there is no evidence to suggest that postmortem damage has a bias toward codon positions (sites $\chi^2 P = .47$; hits $\chi^2 P = .14$).

Regional and Species Specificity

Table 6 presents sample-specific rates of postmortem damage, Π , and saturation, S , for each sample with both HVR1 and COIII data. Six of eight human samples and two of three bear samples demonstrate higher postmortem-damage rates in the HVR1 than in COIII/Cytb, with HVR1 subregions showing 1.1–6.8 times the postmortem damage than corresponding COIII/Cytb regions. Of the three exceptions, two (bear IB221 and human EA) demonstrate HVR1 saturation values of 1, indicating that the postmortem damage has been underestimated. When these two samples are removed, the H_0 of an equal amount of postmortem damage in each region can be rejected (one-way t test $P = .033$; Wilcoxon $P = .038$).

Haplogroup Analysis

Figure 5 shows the relationship between phylogenetic changes and postmortem-damage rates on each branch of the Eurasian HVR1 mitochondrial tree. The analysis is limited because the majority of the samples that we studied are genetically similar northwestern Europeans (Anglo-Saxons and Scandinavians/Vikings). Contemporary indigenous European populations reach up to 60% haplogroup H and 40% haplogroup V (Torroni et al. 1998), and this is mirrored in our findings (M.T.P.G., unpublished data). Consequently, the postmortem damage in the HVR1 is mostly constrained to non-H and non-V substitutions, and unidirectional changes are almost exclusively observed (e.g., postmortem damage at position 16223 will nearly always be constrained to changes from the haplogroup H or haplogroup V cytosine to a 5-hydroxyuracil).

However, where the opportunity for back mutations exists, such as at position 16298 in haplogroup V (changing back to the CRS state), this is observed in only one clone out of the seven specimens. This scenario is marked on the tree. The data permitted a rough estimation of which haplogroups are most likely to arise through postmortem damage, starting from the CRS (table 3). The limiting factors are naturally the number of sites that are needed to change (two sites are less likely to change than one), as well as the site with lowest rate of postmortem damage in a haplogroup. For example, haplogroup I differs from the CRS at sites 16129, 16223, and 16391, which have relative rates of postmortem damage of 12.5, 10.3, and 0, respectively. Although two of these sites are highly susceptible to postmortem damage, site 16391 effectively prevents the misidentification of haplogroup I.

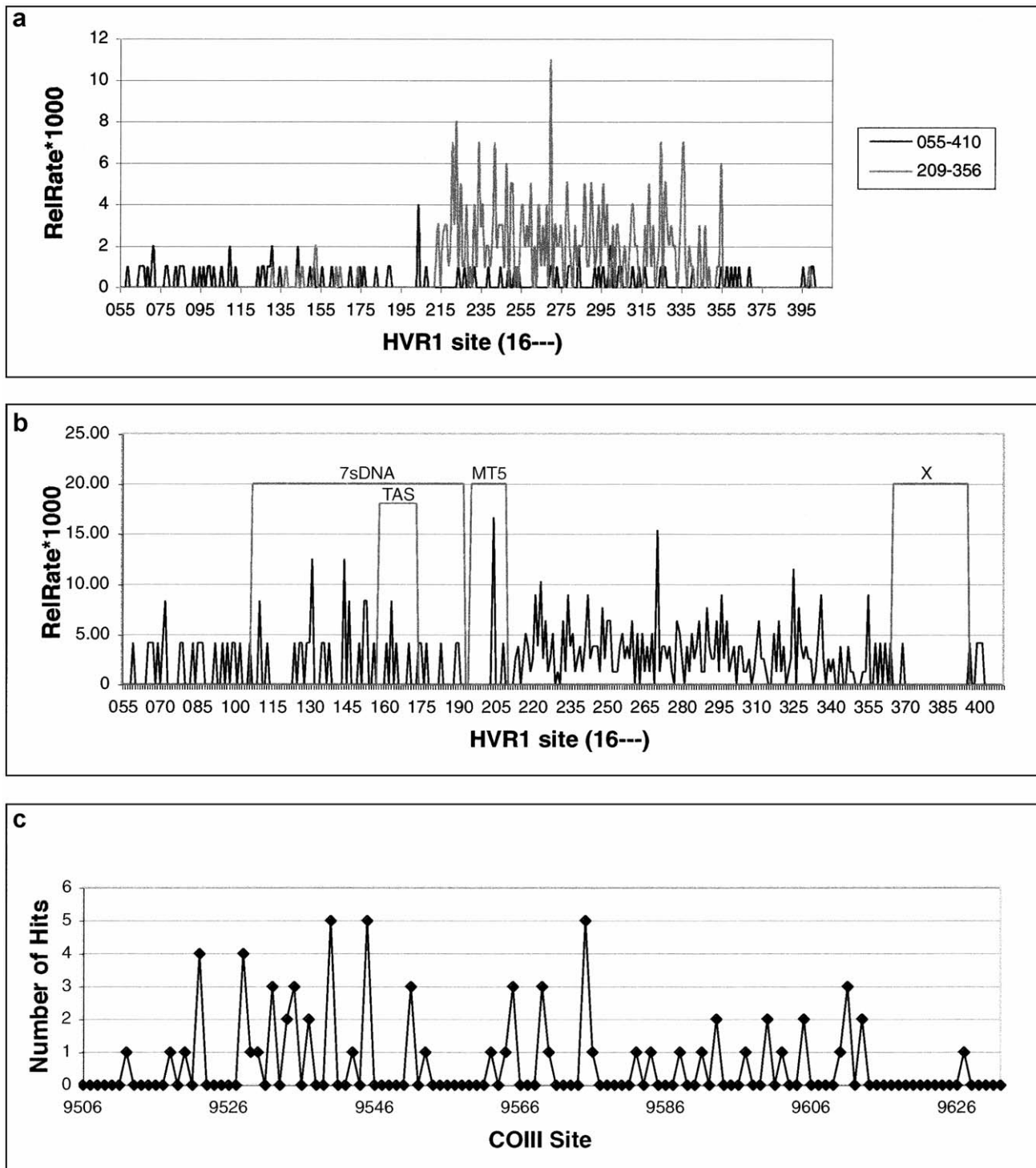


Figure 3 Damage variation across HVR1 and COIII. *a*, Absolute damage measured in hits per site for data sets generated by each PCR primer set across HVR1 (16055–16410). *b*, Relative-rate variation ($\times 100$) across HVR1, calculated as in text. Four areas of interest are marked onto the graph. *c*, Absolute damage across region of COIII (09500–09632). Numbering is with reference to the CRS (Anderson et al. 1981).

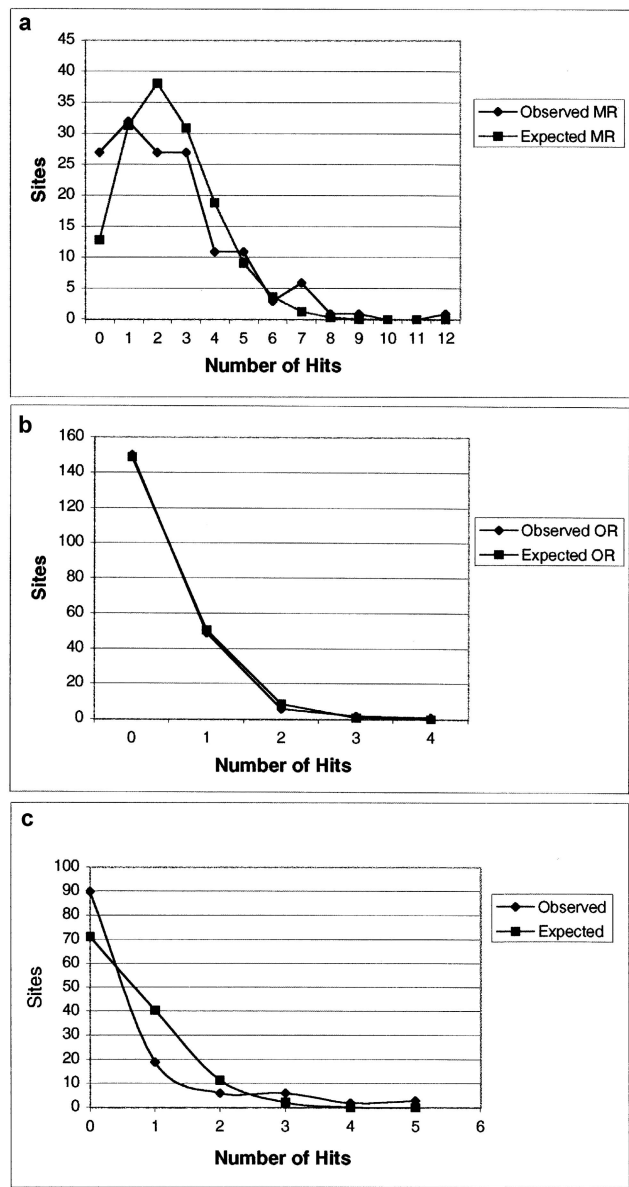


Figure 4 Observed and expected frequency of absolute damage rates for HVR1 MR (a), HVR1 OR (b), and COIII (c) sites. For full details, see text.

Discussion

The sequence results of the present study are consistent with derivation from a series of different specimens containing single aDNA sources. The cloning and UNG experiments, along with the repeat extractions and amplifications, suggest that mtDNA heteroplasmy (Gocke et al. 1998), nuclear copies of mitochondrial genes (Mourier et al. 2001), or polymerase misincorporations (Hansen et al. 2001) are not contributing to the observed variation. The results clearly demonstrate that postmortem DNA

damage is not distributed randomly across the HVR1 or COIII.

Although the HVR1 contains no functional genes, modern population studies have shown that it does not evolve in a random manner, since sequence variation is concentrated at a few sites. Relative rates of mutations have been calculated using a range of methods (Hasegawa et al. 1993; Wakely 1993; Excoffier and Yang 1999; Meyer et al. 1999; Heyer et al. 2001) that are not directly comparable, although striking similarities are apparent even after standardization has reduced the resolution (table 4). Of the 30 sites that can be compared for postmortem-damage and in vivo mutation rates, 15 show very similar rates, and only 6 (sites 16110, 16144, 16148, 16204, 16242, and 16325) completely disagree (i.e., are not observed to mutate in vivo but experience fast postmortem damage). Of these six, at least three have mutation-rate estimates, in the two modern studies, that also dis-

Table 4
Standardized Mutation and Damage Rates

HVR1 Base Position	E99	M99	TG03
16072	2	0	3
16085	3	0	3
16093	4	3	4
16110 ^a	0	0	4
16126	0	4	3
16131 ^a	0	0	3
16144 ^a	0	0	3
16148	3	3	1
16163	4	3	3
16172	3	3	1
16183	3	3	2
16187	4	3	1
16189	4	4	1
16192	4	3	1
16204 ^a	0	0	4
16219	4	3	1
16223	3	4	4
16230	3	4	1
16242 ^a	0	0	3
16262	4	0	3
16270	3	3	4
16278	3	4	2
16290	0	2	3
16293	3	4	1
16294	3	4	2
16298	4	2	4
16309	3	4	1
16319	4	3	2
16325 ^a	0	0	4
16327	4	2	3

NOTE.— Site-specific in vivo mutation rates taken from two previous studies (Excoffier and Yang 1999 [E99]; Meyer et al. 1999 [M99]) were standardized into quartiles and were compared with the standardized postmortem-damage rates from the present study (TG03).
^a Seven sites where major disagreement is observed between rates of occurrence of modern mutations and ancient damage.

Table 5

Statistical Tests on Absolute Number of Sites Changing across the OR and Four Subregions

REGION	BASES	CHANGES ^b	AVERAGE ^c	TWO-SAMPLE <i>t</i> TEST		MANN-WHITNEY NONPARAMETRIC TEST	
				<i>t</i>	<i>P</i>	<i>W</i>	<i>P</i>
OR	208	58	.28				
7sDNA	86	21	.24	−.93	.82	12,435	... ^d
TAS	16	3	.19	.27	.40	20,112	.39
MT5	15	1	.07	1.62	.06	20,396	.10
LDR1	31	1	.03	6.07	.00	19,291.5	.00

^a Size (in bp) of the region.^b Number of damaged sites.^c Average damage per site.^d No *P* value returned by Minitab.

agree, and this may relate to sampling stochasticity or the standardization approach. If so, further sampling may provide evidence for elevated mutation rates at these sites. Other explanations for different rates include the possibility that some form of *in vivo* protection for these sites may be removed or degraded after death.

Structural constraints may also explain the correlation between the postmortem-damage and mutation rates of mtDNA in modern studies. For example, if DNA secondary-structural conformation predisposed particular sites to hydrolytic attack, then these could appear both to be hypermutable in modern populations and to be frequently damaged in postmortem samples. Secondary-structural models of the human HVR1 may provide useful insight but are currently insufficiently detailed. Meyer et al. (1999) remark on the correlation between HVR1 site-specific mutation rates and the three known features of structural interest. A short fragment of heavy-strand DNA, 7sDNA, provides the D-loop with its characteristic triple-stranded feature, extending from a trinucleotide stop codon at position 16104 to at least position 00110 (Doda et al. 1981; Meyer et al. 1999). The stop codon itself exhibits neither postmortem damage nor high mutation rates in data from the present study or previous studies (Excoffier and Yang 1999; Meyer et al. 1999), and 7sDNA itself appears to mutate and receive postmortem damage at average rates. The other two structural regions, TAS (the termination-associated sequence) and the putative control element MT5, have low observed *in vivo* mutation rates, which have previously been suggested to result from functional constraints (Meyer et al. 1999). TAS is located upstream from the trinucleotide stop codon and interacts with sequence-specific binding factors (Doda et al. 1981; Wallace et al. 1995), whereas MT5 is postulated as a protein-binding site (Meyer et al. 1999). Both TAS and MT5 appear to have lower than average postmortem-damage rates, although this cannot be statistically validated at $P < .05$. Certain proteins have been shown to

survive for much longer than DNA (Bada et al. 1999), and it is possible that the low amount of postmortem damage observed in the MT5 region (fig. 3b) is related to the continued binding and protection offered by the putative control element.

The results also identify a previously unrecognized region, LDR1, that possesses significantly reduced levels of postmortem damage. HVR1-mutation-rate studies (Excoffier and Yang 1999; Meyer et al. 1999) do not include all of this region but do characterize low rates of *in vivo* mutation in the 5' segment. The absence of postmortem damage over such a large region (and over 600 clones) is unusual and suggests that the DNA sequence may be protected from hydrolytic damage in some way. One possibility would be a protein-DNA association, similar to that postulated for MT5 (Meyer et al. 1999), and, if further studies verify this observation, then LDR1 may be the first structural feature identified by aDNA. The hypothesis that there is a correlation between function and postmortem-damage rate should be tested further by examining ancient sequences of other CR regions with known binding sites, such as the main regulatory elements of transcription and regulation, or the origin of heavy-strand replication (Chang and Wallace 1985; Wallace et al. 1995).

One of the most interesting findings of the present study is the observation in both humans and bear species that postmortem-damage rates are higher in the HVR1 than in coding regions. Three samples did not show this pattern, two of which (one human and one bear) can be excluded because of high "saturation" values of clone sequence diversity within the HVR1, indicating that the true level of postmortem damage is underestimated; the third anomaly, human sample Tg148, cannot be excluded on this basis, and it is possible that varying amounts of template in the original PCRs may be responsible. Finnilä et al. (2001) note that *in vivo* mutation rates of mitochondrial 3rd codon positions are also lower than those of the HVR1. A plausible explanation is that the secondary-structural conformation of the HVR1 promotes

Table 6**Region-Specific Damage Rates per Sample**

Species, Sample, and Region ^a	Damage Rate (II) ^b	Saturation (S) ^c	Source ^d
<i>H. sapiens</i> :			
76:			
HVR1	1.35	.40	
COIII	.79	.50	
80:			
HVR1	2.82	.86	
COIII	.79	.20	
114:			
HVR1	4.96	.80	
COIII	3.97	.36	
136:			
HVR1	4.81	.41	
COIII	.00	.41	
148:			
HVR1	.68	.18	
COIII	2.91	.30	
149:			
HVR1	5.01	.88	
COIII	4.76	.58	
192:			
HVR1	.82	.40	
COIII	.00	.00	
EA:			
HVR1	.91	1.00	
COIII	1.59	.60	
<i>U. spelaeus</i> TAB15:			
CR	3.4	1.0	Loreille et al. 2001
Cytb	.5	.9	Loreille et al. 2001
<i>U. arctos</i> :			
IB221:			
CR	6.0	1.0	Barnes et al. 2002
12s	9.0	.8	
IB223:			
CR	5.4	.7	Barnes et al. 2002
12s	3.3	.7	

^a Regions given are fragments from the following genes/noncoding regions: COIII, control region (CR), apocytocrome b (Cytb), HVR1, and 12s ribosomal RNA (12s).

^b Sample-specific calculation of damage level (comparable only within regions from one sample).

^c Proportion of clones that vary from each other within one cloned region.

^d Where no source is indicated, data are from the present study.

increased rates of both in vivo mutation and postmortem damage, whereas, in the coding region, either this is lacking or there has been some selection to constrain mutation rates. Structural models of the human HVR1 may provide a useful test of this hypothesis.

It is interesting that the postmortem-damage results do not show the selection against 1st- and 2nd-codon-position mutations seen in coding-region sequences of modern populations. The ancient sequences may actually reflect the real mutation rates of COIII sites; these rates are normally masked by the effects of selection. This pattern shows no evidence for codon bias but does

shows postmortem-damage hotspots, such as the HVR1 sequences.

Problems with Studies on Human aDNA

Current human aDNA research focuses on the identification of mtDNA haplogroups through either restriction digests (e.g., see Merriweather et al. 1994; Kaestle and Glenn Smith 2001) or sequence analysis (e.g., see Hänni et al. 1994; Krings et al. 1997, 1999; Stone and Stoneking 1998; Adcock et al. 2001) and on sexing individuals through systems such as amelogenin (e.g., see Faerman et al. 1995; Fily et al. 1998; Vernesi et al. 1999). The sexing and restriction-digest studies lack sufficient resolution to allow the assessment of the role of postmortem damage or contamination, so we can only analyze research that uses sequence data to define—or even determine—new haplogroups, such as studies of Neanderthal or archaic human samples (Krings et al. 1997, 1999; Ovchinnikov et al. 2000; Adcock et al. 2001).

The data in the present study have been generated from a small set of northwestern European samples, predominantly of haplogroups H and V. Consequently, the mtDNA tree (fig. 5) has been modified to demonstrate the potential haplogroup alteration caused by damage to these haplogroups only. However, figure 5 demonstrates the ease with which phylogenetic misidentification can occur if postmortem damage is not detected in sequences from any haplogroup. For example, it is not inconceivable that the Amerindian haplogroup A sequence could result from a haplogroup H Viking sample. We suggest that postmortem damage may explain many unusual results obtained from ancient human remains when appropriate techniques were not followed (e.g., the 60,000-year-old “Mungo man” sequences; Adcock et al. 2001). Postmortem damage will also complicate population genetic analyses of ancient humans, and detailed cloning will be needed to avoid overestimation of heterogeneity and population expansion sizes (Lundstrom et al. 1992; Aris-Brisou and Excoffier 1996).

For human aDNA research to progress, it will be necessary to rigorously enforce aDNA guidelines (Cooper and Poinar 2000). As real-time PCR becomes more routinely available, the initial template copy number should be quantified in samples, and detailed cloning experiments should become a requirement. Samples with low template number should be replicated and cloned at least once, since even highly damaged sites, such as 16270, are unlikely to predominate in two independent PCRs. However, the simplest procedure appears to be a comparison of sequences before and after enzymatic treatment with UNG (Pääbo 1989; Hofreiter et al. 2001), EndoIV (Pääbo 1989), AAG (Lau et al. 2000), or T4 with Pol 1 (Pusch et al. 1998; Di Bernardo et al. 2002), to reduce the number of templates with postmortem-damaged bases and the

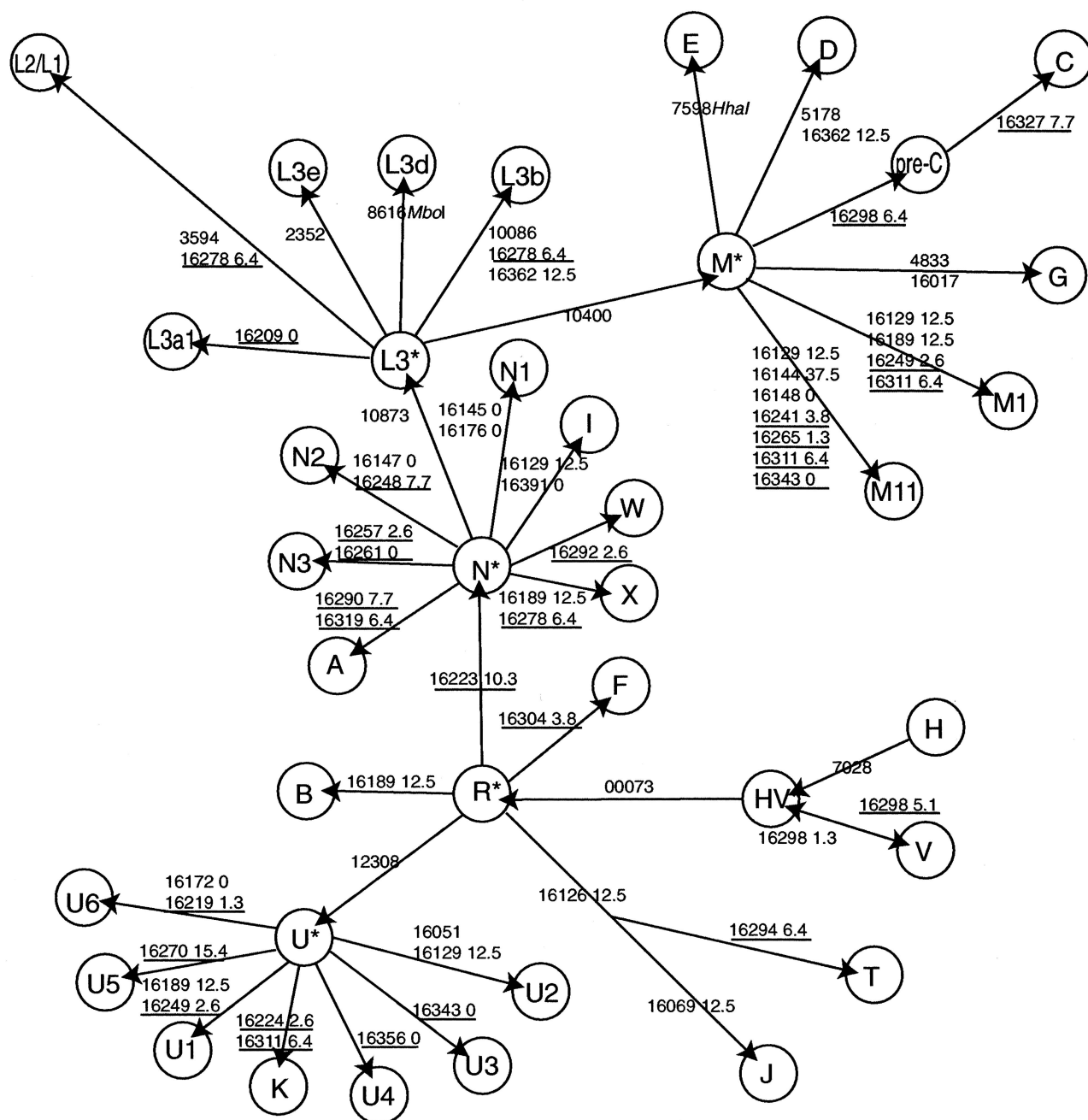


Figure 5 Modification of mitochondrial HVR1 haplogroup tree (reprinted, with permission, from Richards and Macaulay 2000) to demonstrate potential for haplogroup misidentification. From a starting sample of haplogroups H or V, arrows indicate the direction along the tree a sample can move because of hydrolytic deamination (or real singleton substitution) of C→T or A→G. Site numbers are with reference to the CRS (Anderson et al. 1981). Relative rates are calculated for the MR (underlined) and OR. For full details, see text.

resulting possibility of misidentification. Such treatment is expected to reduce the starting-template copy number, providing a check for authenticity but also limiting the number of samples suitable for study. A variety of other nongenetic factors should also be considered, such as depositional environment (Höss et al. 1996; Nielsen-Marsh 2000) and microbial content (Burger et al. 1999), amino acid racemization (Poinar et al. 1996; Poinar and Stanekiewicz 1999) and composition (Bada 1999), and gas chromatography/mass spectroscopy-measured levels of hydantoins (Höss et al. 1996). These issues are now well known, and the heavy burden of proof associated with aDNA research demands that new studies address them.

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